

PAJUNK® GmbH Medizintechnologie • Karl-Hall-Str. 1 • 78187 Geisingen, Germany

Geisingen, 2019-09-02

# **Urgent Field Safety Notice**

## **RECALL OF CERTAIN BATCHES**

of

## **SPROTTE®** lumbal with Introducer

\_\_\_\_\_

#### **SENDER:**

PAJUNK® GmbH Medizintechnologie Karl-Hall-Str. 1 78187 Geisingen

#### **RECIPIENT:**

Xxx

Xxx

XXX

#### **IDENTIFICATION OF AFFECTED DEVICES:**

Trade Name: SPROTTE® Lumbar with Introducer

Item number(s): see ATTACHMENT I BATCH 1240 - 1313

#### **Dear valued Customer,**

PAJUNK® GmbH Medizintechnologie has internally identified a packaging problem that may affect certain batches of the SPROTTE® lumbar puncture cannulas with introducer listed in Attachment 1.

The cannulas type "SPROTTE®" are used for diagnostic lumbar puncture / puncture of the spinal space for CSF collection.

 $info@pajunk.com \cdot www.pajunk.com\\$ 



This letter is meant to inform you about the problem, explain the measures you have to take and the actions that PAJUNK has in place to address the issue.

### Affected products

The complete list of affected products including item number is attached to this letter (Attachment 1).

#### Description of product problem

PAJUNK® GmbH Medizintechnologie received information about a problem which has occurred in the packaging sealing process during the manufacturing of certain products.

Due to this problem, PAJUNK® GmbH Medizintechnologie cannot guarantee with sufficient certainty that the sterilized medical devices to which this safety measure applies remain reliably sterile during their defined storage and shelf life.

The problem could be identified and limited to the products listed in the attachment. To avert potential hazards, PAJUNK® GmbH Medizintechnologie has decided to recall the affected products.

## Description of the potential consequences to patients:

In the case of failure to comply with this customer information there is a risk of using a nonsterile product on the patient.

#### Action to be taken by the recipient

- 1. Identify the affected products (per Attachment 1) and quarantine!
- 2. Do not use any of the affected products!
- 3. Please fill in and return the attached reply form (Attachment 2) accompanied by the affected products to your contact point at PAJUNK®/ your distributor of PAJUNK®devices.

## Further actions planned by PAJUNK® GmbH Medizintechnologie

PAJUNK® GmbH Medizintechnologie has reviewed the packaging sealing process, taken corrective action and will implement preventive actions to ensure the highest level of product safety and quality.

PAJUNK® GmbH Medizintechnologie will replace the returned devices subject to this recall free of charge and without any additional order within 2 – 5 weeks.

Commerzbank AG



## **Transmission of this Field Safety Notice**

This notice needs to be passed on all those who need to be aware within your organisation. Please transfer this notice to any organisation on which this action has an impact or inform below mentioned contact person about third parties where the affected products have been transferred

Please retain this information at least until the measure has been completed by PAJUNK® GmbH Medizintechnologie. Please maintain awareness on this notice and resulting action for an appropriate period to ensure effectiveness of the corrective action.

Please report all device-related incidents to the manufacturer, distributor or local representative, and the national Competent Authority if appropriate.

Your national Competent Authority, the Agencija za lijekove i medicinske proizvode, Ksaverska cesta 4, 10000 Zagreb, has received a copy of this "Urgent safety information: RECALL of a Medical Device".

#### Contact person logstics / customer service:

Ms. Nilüfer Sen PAJUNK® GmbH Medizintechnologie Karl-Hall-Strasse 1 78187 Geisingen Baden-Wuerttemberg, Germany Fon +49(0)7704-9291 ext. 647 Fax +49(0)7704-9291 ext. 600 Niluefer.sen@pajunk.com

## Contact person Regulatory Affairs / Safety Officer:

Christian G. H. Quass

Director Regulatory Affairs & Safety Officer for Medical Devices

PAJUNK® GmbH Medizintechnologie

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## **Attachment 1**

# List of affected products

Item Number	LOT
321151-30C SPROTTE Lumbal	1240 to 1313
331151-31B SPROTTE Lumbal	1240 to 1313

 $info@pajunk.com \cdot www.pajunk.com\\$ 



# Attachment 2 Reply Form

Please return this form together with the original letter within 5 days of receipt of the urgent safety information by fax, letter or e-mail attachment to the person named in the cover letter or to **sibe@pajunk.com** 

Recipient:	Sender [stamp/physical address of institution]
PAJUNK® GmbH Medizintechnologie -Sicherheitsbeauftragter- Karl-Hall-Strasse 1	
78187 Geisingen	
We hereby confirm receipt of the aforemention	ned urgent safety information.
We have identified affected devices (If multiple batches or multiple article numbers are involved you kindly submit a detailed breakdown.)	s in our institution. lved, PAJUNK® GmbH Medizintechnologie requests that
Number of devices/ individual packs that we a	re immediately returning:
Number of affected devices that have already	been used on patients to date:
SIGNATURE AREA	
Name/ position [BLOCK LETTERS]	Date/ signature